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(71) Applicant: IDEXX LABORATORIES, INC. [US/UI] Idexx Drive, Westbrook, ME 04092 (US).	JS]; Or	ne		ML, MR, NE, SN		
(72) Inventors: MERMER, Brion; 2 Linden Court, Cur ME 04092 (US). HARRIS, Rachel, A.; 254 Buzz Dayton, ME 04005 (US). SIEFRING, Ann, E.; Farm Road, Salmouth, ME 04105 (US).	ell Roa	1 ,		sternational searc pt of that report.	ch report and to be republished	
(74) Agents: LITHGOW, Timothy, J. et al.; Lyon & Lyo First Interstate World Center, Suite 4700, 633 W Street, Los Angeles, CA 90071-2066 (US).						
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2. WALKING TO CLONE EXON 3	(IgE P	PCR 1)			
				(CANINE	PRIMER 1)	
	NESTE	D PCR 2	-		1	
				(CANINE	PRIMER 2)	
	(SELEC	TIVELY	AMPLIFY IgE P	RODUCT OF PCI	R 1)	

(57) Abstract

Recombinant DNA molecules encoding complete canine IgE heavy chain constant region and the DNA sequence for all six exons of canine IgE are disclosed. The canine IgE heavy chain constant regin DNA sequence was isolated using a nested walking procedure that is also disclosed. Also disclosed are peptides encoded by said sequences, including recombinant canine IgE heavy chain peptides produced by prokaryotic or eukaryotic cells. Such peptides are used in methods to treat the manifestation of allergy in dogs. Disclosed are antibodies that bind to neptides disclosed herein, as well as such antibodies for use to treat the manifestations of allergy in dogs.

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DESCRIPTION

Nucleotides and Peptides Corresponding to the
Canine Ige Heavy Chain Constant Region and Related Methods

Cross-reference to Related Applications

This application is a continuation-in-part of U.S. application serial number 08/601,197 filed 14 February 1996 from which priority is claimed; this application is incorporated by reference herein.

Technical Field of the Invention

This application relates to the analysis and manipulation of genetic material. More specifically, the invention relates to recombinant DNA molecules encoding canine immunoglobulin proteins.

Background of the Invention

It is estimated that up to 30% of the canine population suffers from allergies and allergy-related skin disorders. Specifically, atopy (allergic dermatitis) has been estimated to affect between 3 and 15% of the entire population. The substances most likely to cause an allergic reaction vary from species to species. Common canine allergens include fleas, pollens, molds and dust. Given the prevalence of allergies in dogs, it has been desired to develop methods to properly diagnose and treat allergies in dogs.

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Immunoglobulin E (hereinafter IgE) is a type of antibody that is understood to be an important component mediating allergic responses, including immediate hypersensi-tivity. IgE molecules bind to mast 5 cells and basophils by Fc receptors on the cells. such cell-bound IgE antibodies bind to an allergen, the allergen cross-links with other IqE antibodies on the cell This cross-linking mediates Type I immediate surface. hypersensitivity reactions by causing release of 10 histamines and other molecules that produce symptoms associated with allergy.

Detection and quantitation of IgE antibodies is important in the diagnosis of Type 1 (immediate-type) hypersensitivity disease. Currently, it is understood 15 that canine IgE is detected by several commercially available in vitro allergy tests. Such tests were designed to detect canine IqE in a sample, by contacting the sample with immobilized allergen, then any IgE bound to the immobilized allergen is detected with a polyclonal 20 antibody understood to react with IgE. It is not clear, however, that the polyclonal antibodies react exclusively with IgE since the immunogens used to prepare the polyclonal antibodies were partially purified, native, glycosylated immunoglobulins. Polyclonal antibodies to 25 such immunogens are likely to detect non-IgE immunoglobulins with anti-allergen specificity, since the other immunoglobulins likely contaminated the immunogen. antibodies addition, directed antibody against

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glycosylations are likely to cross-react with glycosylations on IgE and non-IgE immunoglobulins.

The sequences for the genes encoding human and murine IgE heavy chain constant region are known (For example, Ishida et al., "The Nucleotide Sequence of the Mouse Immunoglobulin E Gene: Comparison with the Human Epsilon Gene Sequence", EMBO Journal 1:1117-1123 (1982). human and murine genes possess 60% homology within exons, 45-50% homology within introns, with insertions and deletions. Subsequent to the present invention, Patel et al. published the nucleotide and predicted amino acid sequence for exons 1-4 of the heavy chain constant region of canine IgE in the article "Sequence of the Dog Immunoglobulin Alpha and Epsilon 15 Constant Region Genes, " Immunogenetics 41:282-286 (22 March 1995). The complete sequence of the canine IgE heavy chain constant region, with membrane bound portions encoded by exons 5 and 6, is disclosed for the first time herein.

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Summary of the Invention

The present invention relates to recombinant DNA molecules, and conservative variants thereof, that encode elements of the structural gene for canine IgE. The complete nucleotide sequence that encodes the canine IgE heavy chain constant region (epsilon) is disclosed for the first time herein.

In one aspect of the present invention a recombinant DNA molecule, and conservative variants thereof,

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comprising DNA sequence of exons one, two, three, four, five, or six of the canine IgE heavy chain constant region molecule is set forth.

In another aspect of the present invention,

5 recombinant vectors, such as expression vectors,

comprising a DNA sequence encoding canine IgE heavy chain

constant region or components thereof are set forth.

In another aspect of the present invention, polypeptides produced using a recombinant expression vector containing a canine IgE heavy chain DNA sequence are set forth.

In another aspect of the invention, compositions comprising a nucleotide sequence that is antisense to a DNA sequence encoding a component of canine IgE heavy chain is set forth; the component can be the full IgE heavy chain.

Disclosed is a cloning vector comprising a DNA sequence of canine IgE genomic material; a recombinant cell line comprising a DNA sequence of canine IgE genomic material; a method for producing a polypeptide comprising a step of expressing peptide comprising an amino acid sequence encoded by a cloning vector comprising canine IgE genomic material; a cell that expresses a recombinant polypeptide encoded by a DNA sequence (or conservative variants thereof) that encodes canine IgE heavy chain constant region or components thereof.

Accordingly, an object of the present invention is to provide the full DNA sequence for the canine IgE constant region molecule.

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Brief Description of The Drawings

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Figure 1 is a schematic outline of the nested walking procedure used for the initial cloning of the exons 1-4.

Figure 2 depicts a comparison of cysteine conservation between a portion (exons 1-4) of the sequence encoding canine IgE heavy chain isolated herein (Seq 3 in Fig. 2), compared to portions of murine (Seq 2 in Fig. 2) and human (Seq 1 in Fig. 2) sequences encoding IgE heavy chain constant regions; (*) indicates cysteine, the arrow indicates the onset of canine exon 4.

Figure 3 shows the DNA sequence and predicted amino acid sequence of exons 1-4, and surrounding noncoding DNA of the sequence encoding canine IgE heavy chain constant region. N = unknown nucleotide; Xaa = unknown amino acid.

Figure 4 shows the DNA sequence and predicted amino acid sequence of exons 5 and 6, and surrounding noncoding DNA, of the canine DNA sequence encoding the IgE heavy chain constant region. The probable beginning of exons 5 and 6 are shown.

Figure 5 shows the nucleic acid and amino acid sequences for the signal (secretion) sequence utilized to obtain expression of selected nucleic acid sequences of canine IgE DNA in eukaryotic cells, ligated with canine IgE heavy chain exons 2-4. Upon creation of a recombinant nucleic acid sequence containing the signal sequence, recombinant peptides having functions of native canine IgE were expressed in eukaryotic cells without, however, the need for nucleic acid sequences encoding the entirety of

IgE heavy chain constant and variable regions as well as the light chains.

Detailed Description of The Invention

5 <u>Definitions</u>

cDNA clone: A duplex DNA sequence representing an RNA, carried in a cloning vector.

Cloning: The selection and propagation of a single DNA species.

Cloning Vector: A plasmid, phage DNA or other DNA sequences, able to replicate in a host cell and capable of carrying exogenously added DNA sequence for purposes of amplification or expression of the added DNA sequence.

Codon: A triplet of nucleotides that represents an amino acid or termination signal.

DNA Sequence: A linear series of nucleotides connected one to the other by phosphodiester bonds between the 3' and 5' carbons of adjacent pentoses.

Expression: The process undergone by a structural 20 gene to produce a polypeptide. It is a combination of transcription and translation.

Expression Control Sequence: A DNA sequence of nucleotides that controls and regulates expression of structural genes when operatively linked to those genes.

25 Exon: A contiguous region of DNA encoding a portion of a polypeptide. Reference to any exon, e.g. "DNA sequence of exon 6", refers to the complete exon or any portion thereof.

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Genome: The entire DNA of a substance. It includes inter alia the structural genes encoding for the polypeptides of the substance, as well as operator, promotor and ribosome binding and interaction sequences such as the Shine-Dalgarno sequences.

Nucleotide: A monomeric unit of DNA or RNA consisting of a sugar moiety (pentose), a phosphate, and a nitrogenous heterocyclic base. The four DNA bases are adenine ("A"), guanine ("G"), cytosine ("C") and thymine ("T"). The four RNA bases are A, G, C and uracil ("U"). A and G are purines, and C, T, and U are pyrimidines.

Phage or Bacteriophage: Bacterial virus many of which include DNA sequences encapsidated in a protein envelope or coat ("capsid").

15 Plasmid: An autonomous self-replicating extrachromosomal circular DNA.

Polymerase Chain Reaction (PCR): A method of amplifying a target DNA sequence contained in a mixture of DNA sequences, by using oligonucleotide primers that flank the target DNA sequence for repeated cycles of DNA synthesis of the target DNA sequence.

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Polypeptide: A linear series of amino acids connected one to the other by peptide bonds between the α -amino and carboxy groups of adjacent amino acids.

Reading Frame: The grouping of codons during translation of mRNA into amino acid sequences. For example, the sequence GCTGGTGTAAG may be translated in three reading frames or phases, each of which affords a different amino acid sequence:

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GCT GGT TGT AAG-Ala-Gly-Cys-Lys
G CTG GTT GTA AG-Leu-Val-Val
GC TGG TTG TAA A-Trp-Leu(STOP).

Recombinant DNA Molecule: A hybrid DNA sequence comprising at least two nucleotide sequences, the first sequence not normally being found together in nature with the second.

Structural Gene: A DNA sequence which encodes through

10 its template or messenger RNA ("mRNA") a sequence of amino
acids characteristic of a specific polypeptide.

Transcription: Synthesis of RNA on a DNA template.

Translation: Synthesis of protein on the mRNA template.

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Initial Cloning of Canine DNA Corresponding to Exons 1-4 of the IgE Heavy Chain Constant Region

fragment from a mixture of recombinant DNA clones is to
use a DNA probe derived from a known target DNA sequence.
Prior to the present invention, however, canine IgE DNA
was not available. In such situations a number of methods
in the art are used to isolate a gene from one species,
based on homology with the corresponding gene of known
sequence from another species. Typically, DNA fragments
derived from homologous genes from more distantly related
species, show some small regions of greater homology
scattered among regions of much lower homology. The more
homologous segments form the basis for interspecies

identification with such probes. Therefore, DNA derived from the IgE gene from at least one other species is used. In order for this method to be practical, however, sufficient homology must be found between the target DNA and the probe. If homology is not sufficient, it will not be possible to distinguish binding of probe to target from nonspecific binding. It was known that there was a low degree of homology between mouse and human IgE heavy chain constant region genes. Accordingly, a low degree of homology between canine IgE heavy chain constant region genes and the genes of these other species was also expected; this low degree of homology made the use of prior art methods difficult.

Accordingly, in an effort to clone the gene encoding the canine IgE heavy chain constant region (epsilon), a 15 (PCR) polymerase chain reaction method based on A PCR-based "walking" amplification was developed. approach was used on canine genomic DNA isolated from whole blood (Promega, Inc.). Although less direct than 20 having a known probe sequence, PCR using oligonucleotides has several advantages over direct DNA probe methods. First, it is more practical with PCR to employ a spectrum of different physical parameters such as temperature in order to allow specific binding to target Second, the size of the resulting PCR product shows 25 DNA. that not only does a single gene contain homology with two oligonucleotide primers, but their relative locations are as predicted. The combination of these factors makes it

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more likely that PCR will successfully locate DNA fragments based on partially homologous oligonucleotides.

The PCR walking approach does not guarantee that interspecies homology will effectively allow isolation of homologous DNA fragments. There is reliance on primers derived from other species; thus, if the homology with such other species is insufficient, no target fragments will be obtained.

In the present cloning of DNA encoding canine IgE 10 heavy chain constant region, a step was introduced in an effort to insure that authentic canine IgE-encoding fragments were isolated by PCR. The step was based on the concepts of "genomic walking" and of nested PCR. Genomic walking refers to isolation of a DNA fragment prepared from the end of a starting DNA, and its subsequent use as a probe to isolate an overlapping cloned DNA fragment Nested PCR is a which also contains the probe region. technique designed to increase the specificity of PCR amplification. With nested PCR, the products of an 20 initial PCR are reamplified with a second set of primers selected from within the specific predicted product. Only the specific product, and not any unrelated products, should be amplified in the second reaction. Nested PCR is commonly used when the target DNA is present at low levels in a starting sample likely to contain many non-specific 25 targets. In such situations, a relatively high contribution of non-specific products in the final set of products is expected. A second, nested PCR, should greatly increase the proportion of specific products.

the typical nested PCR situation, however, the sequence of the target is known, and only primers which match that target are used.

To adapt the genomic walking concept to PCR, and to use the method to isolate fragments based on interspecies homology, an oligonucleotide from one end of a cloned fragment was used as a primer; a second primer was determined based on the possibility of interspecies homology (rather than being derived from a known sequence from the same species as with usual nested PCR). To enhance the specificity of the present approach, a second PCR reaction was performed. The products of the first reaction were amplified substituting a different primer obtained from within the starting cloned fragment. Only one of the primers in the second PCR was specific for the authentic product, unlike the common case with nested PCR where all primers are specific for a known product.

The PCR-based "walking" approach employed herein is depicted in Figure 1. This walking approach was used to clone, consecutively, genomic fragments containing copies of exons 4, 3, 2, and 1, including the intervening sequences separating them.

This PCR-based method was carried out as follows:

One end of the target gene (exon 4 in this case) was

cloned by PCR with a primer pair selected from either a

murine or a human IgE heavy chain constant region-encoding

sequence. These PCR primers were selected from regions

which demonstrated relatively high sequence conservation

between the murine and human sequences.

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An initial fragment of canine exon 4 was obtained after cloning and sequencing products from a variety of different PCR reactions with different primer pairs. The fragment was determined to be canine exon 4 based on comparison of DNA and amino acid sequences between murine and human exon 4. Although the degree of homology was low, the pattern of conserved residues was similar among the three sequences (Fig. 2). Once the initial canine fragment of exon 4 was sequenced, additional clones containing adjacent DNA were isolated by the nested PCR-based "walking" approach.

To obtain clones of adjacent exon 3 (and intron 3), a PCR (referred to as PCR1) was performed with one primer selected from the mouse or human exon 3 sequences and with a second primer from within the cloned canine exon 4 sequence. In PCR1 at least one primer would be a match with the target, the primer derived from the authentic canine exon 4 sequence. Since one of the primers was likely not to be a precise match, the products of the PCR reaction include a set of "false" products detected as an unresolved "smear", presumably containing the authentic product.

To amplify the authentic product but not the false products and to extend the cloned DNA into the adjacent exon 3 (and intron 3), a second nested PCR reaction with a different canine primer was performed using the products of the PCR1 as the targets. In this particular case, the primers included the non-canine exon 3 primer which was used in PCR1, and a second authentic canine primer from

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within exon 4. Among the products of the first reaction, only the authentic IgE heavy chain constant region product contained exon 4 and therefore was amplified in the second PCR. The fragment resulting was cloned and sequenced.

This nested "walking" procedure was used to clone, consecutively, genomic fragments containing copies of exons 4, 3, 2 and 1, including the intervening sequences separating them. Nested PCR reactions which involved one "authentic" canine primer were performed under conditions of higher stringency, so that fewer DNA sequences were modified and so that the sequences modified more closely correspond to the primers.

The PCR products were cloned in the plasmid pUC19 (Boehringer-Mannheim), and the DNA sequence was determined. Sequence comparison with the human and mouse sequences indicated that the product represented canine IgE heavy chain constant region, since even though the homology was low, when there was a sequence match between two of the species, the match often tended to occur in the genes of each of mouse, human and dog. For example, a comparison of cysteine conservation, an important amino acid in immunoglobulin structure, is depicted in Figure 2.

The PCR primers used herein are indicated in Table 1.

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Table 1

IgE heavy
chain constant
region exon,
plasmid
(plasmid

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synonyms) PCR PCR primers

	,		
10	exon 4 191gE (191gE-4)		TGATCCAGAACTTCATGCCTGAGGA AGGCGACTGAAGATGAAGAAGCC
	exon 3 19IgE3/4	PCR 1	CCTACCTAAGCCGGCCCAGCCCGTTCGACCTGTT AGGCGACTGAAGATGAAGAAGCC
		Nested PCR 2	CCTACCTAAGCCGGCCCAGCCCGTTCGACCTGTT CCGTGGTGGTGTACTGGTCT
15	exon 2 19ce23	PCR 1	AGAAGCACTGGCTGTCAGACCGCACCTACACCTG TAGACGTGACTGTGATCGTCC
		Nested PCR 2	AGAAGCACTGGCTGTCAGACCGCACCTACACCTG CCGGTACCAGGTCAGGT
	exon 1 19ce12A-1	PCR 1	TGACTCTGGGCTGCCTGGCCACGGGCTACTCCC GGCCATACCTGAGCACTTGCG
		Nested PCR 2	TGACTCTGGGCTGCCTGGCCACGGGCTACTCCC CTGAGCACTTGCGAGCCTCAT

It will be appreciated by one skilled in the art that primers other than those set forth here are also effective in the isolation and amplification DNA encoding canine IgE heavy chain constant region in accordance with the methodology disclosed herein.

It will be appreciated by one of ordinary skill in the art that a wide variety of host/cloning vector combinations are usefully employed in cloning the DNA isolated as abov . For example, useful cloning vehicles/vectors have included pMALc2 (New England Biolabs), and pGST (Pharmacia) to express canine IgE

fragments in bacteria; the canine IgE DNA has been propagated in pUC19 (Boehringer-Mannheim) and λZAP (Stratagene). Cloning vehicles comprise various known bacterial plasmids such as pBR322, other E. coli plasmids 5 and their derivatives and wider host range plasmids such as RP4; phage DNA such as the numerous derivatives of phage λ , e.g., NB989; and, vectors derived from a combination of plasmid and phage DNA, such as plasmids which have been modified to employ phage DNA expression 10 control sequences. Vectors which replicate in eukaryotic cells can also be used. Useful hosts comprise bacterial hosts such as E. coli strains X1776, X2282, HB101 and MRC1; strains of Pseudomonas, Bacillus subtilis and other bacilli; yeasts and other fungi; animal or plant hosts 15 such as animal or plant cells in culture; and other hosts. As appreciated by one of ordinary skill in the art, not all hosts are equally efficient. The particular selection of host-cloning vehicle combination may be made by those of skill in the art after due consideration of the 20 principles set forth herein without departing from the scope of this invention.

Isolation of Genomic Clones Containing Exons 1-6 of Canine IgE Heavy Chain Constant Region DNA

In order to obtain the complete gene for the canine IgE heavy chain constant region, a canine genomic DNA library was prepared and screened using the λ ZAP vector. Genomic DNA was digested with the restriction endonuclease BamHI. A probe was prepared from canine exon 1, cloned as

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described herein; this probe was used to screen the library. A 7 kB canine genomic fragment, clone ce7, was isolated and characterized; the plasmid pBKce7 contained the genomic material of subclone pBKce7; pBKce7 was constructed from the λZAP clone by in vivo excision. DNA sequencing indicated that clone ce7 included exons 1, 2, 3 and part of exon 4 extending through a BamHI site near the end of exon 4. It was determined that BamHI had digested near the end of exon 4 locus based on comparisons with murine and human exon 4, and the absence of a termination codon.

In order to isolate a clone containing the remainder of the canine IgE heavy chain constant region locus, a second genomic library was constructed with genomic DNA after digestion with restriction endonuclease HindIII and the library was screened with the probe prepared from exon 1 which was used in the isolation of the clone ce7. Clone ce5, which resulted, was a 5 kB fragment which contained exons 1, 2, 3, 4, 5, and 6, and a large intron separating 20 exons 5 and 6 from the others; plasmid pBKce5 which was constructed by in vivo excision contained the genomic material of clone ce5. Exons 5 and 6 are believed to encode additional amino acid residues specifically found in the cell-associated form of IgE but not in the secreted 25 form.

Figure 3 shows the DNA sequence and predicted amino acid sequence of exons 1-4 and surrounding noncoding DNA of canine IgE heavy chain constant region. As discussed in greater detail herein, the exon boundaries were

determined based on sequencing cloned cDNA copies of mRNA isolated from canine lymphocytes as discussed herein. The exon boundaries are represented in Fig. 3 (SEQ ID NO: 1) as follows: the DNA sequence of exon 1, nucleotides 167-448; the DNA sequence of exon 2, nucleotides 608-931; the DNA sequence of exon 3, nucleotides 1024-1344; and, the DNA sequence of exon 4, nucleotides 1419-1742. Underlined nucleotides 1835-1840 show the polyadenylation signal for the secreted form.

The complete DNA sequence of the locus containing exons 5 and 6 is indicated in Figure 4. Figure 4 also shows the predicted amino acid sequence of exons 5 and 6, and surrounding noncoding DNA near the canine IgE DNA The beginnings of exons 5 and 6 as presently understood are shown in Fig. 4; the beginning sites were based on comparisons with human sequences, and not on cDNA sequences; the end of each exon was determined based on general sequence motifs known to those of skill in the art to be found at the ends of exons. The precise limits of the exons is known upon examination of the mRNA with the spliced together. The exon boundaries, determined, are depicted in Fig. 4 (SEQ ID NO: 2) as follows: the DNA sequence of exon 5, nucleotides 82-216; and, the DNA sequence of exon 6, nucleotides 316-390.

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Preparation of Canine Genomic DNA

A canine genomic DNA fragment comprising IgE heavy chain epsilon exons, or components thereof, is available is deposited with the ATCC as Accession Numbers (not yet

assigned). Alternatively, a canine genomic DNA fragment comprising IgE epsilon exons, or components thereof, is obtained by isolation of the 5 kilobase HindIII fragment referred to herein as ce5, or portions thereof. To isolate canine genomic DNA corresponding to the sequence of ce5, a library of 5 kilobase fragments is prepared from canine genomic DNA digested with restriction endonuclease HindIII; digestion with this endonuclease was determined to contain complete canine heavy chain constant region exons 1-6 in the section "Isolation of Genomic Clones 10 Containing Exons 1-6 of Canine IgE Heavy Chain Constant Region DNA" herein. The library is screened with a DNA To prepare the probe, PCR is performed with canine genomic DNA as target, with primers constructed in accordance with standard methodologies to amplify a 15 fragment between coordinates 251 and 444 as defined in Fig. 3(SEQ ID No. 1). The sequence of HindIII digested DNA identified with the probe is isolated.

20 Characterization of Messenger RNA for Canine IgE Heavy Chain Constant Region

As indicated herein, sequence comparisons using the deduced amino acid sequence of the cloned DNA showed patterns of conservation consistent with a determination that the canine IgE Heavy chain epsilon sequence was isolated. However, it was possible that the cloned DNA represented a nonfunctional canine gene. To rule out the possibility that the cloned DNA was for a nonfunctional gene, analysis for the presence of mRNA copies of the gene

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in canine lymphocytes was performed. To determine whether lymphocytes contained such mRNA copies, cDNA cloning and Purified mRNA was used. from sequencing lymphocytes was converted into DNA and PCR was performed 5 with primers based on the cloned genomic sequence. protocol was carried out in accordance with the principle that isolation of PCR products whose sequence matched the canine genomic exon sequences would, therefore, indicate that the cloned gene was expressed in dog lymphocytes. Canine lymphocytes were isolated from canine blood and mRNA was purified using standard procedures. The mRNA was converted to double stranded cDNA by reverse transcription and PCR amplification with primers based on the genomic DNA sequence, and cloned into pUC19. Several cDNA clones 15 were identified and sequenced. Such cDNA cloning of mRNA and subsequent sequencing was carried out for several cDNA Sequencing of these clones was performed. clones. DNA sequence of the cDNA clones was as expected for transcripts of the genomic clone described above, confirming that the clone represented an active IgE heavy chain The cDNA sequence also allowed assignment of the exon boundaries for exons 1, 2, 3, and 4, as indicated in Figure 4.

25 Expression of Recombinant IgE Heavy Chain Constant Region Proteins in E. coli.

A portion the DNA sequence encoding exon 3 was expressed as a recombinant fusion protein in *E. coli* by use of a cloning vector (in this context an expression

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vector). The recombinant vector was a derivative of the commercially available plasmid pEX3 (Boehringer Mannheim). The plasmid pEX3 contained the bacteriophage lambda promoter/operator sequence and was designed to produce a fusion protein with β-galactosidase, which was encoded by the lacZ gene derived from E. coli with pEX3; expression was regulated by a temperature shift in the presence of the temperature sensitive cI857 repressor.

The pEX3 expression vector was modified by the deletion of 1885 nucleotides between the restriction enzyme sites EcoRV and SmaI in the β -galactosidase coding region. This deletion resulted in a deletion of about two-thirds of the β -galactosidase coding region, with and left sufficient DNA to encode for approximately 400 amino acids.

A portion of exon 3, encoding sixty-four amino acids, was ligated into the EcoRV/SmaI site of the modified pEX3 plasmid. The portion of exon 3 that was inserted corresponded to nucleotides 1081-1272 (as depicted, e.g., in Fig. 3). The IgE DNA was inserted so that the reading frame of the encoded protein was in frame with the reading frame of the coding region for β-galactosidase, whereby a fusion protein between canine IgE constant region peptide and β-galactosidase was produced. The IgE fragment in plasmid 19IgE3/4 was oriented within vector pUC19 such that this translational fusion was generated by ligation of the 0.22kB EcoRI/SalI fragment from plasmid 19IgE3/4 with the 6.6kB EcoRI/SalI fragment from the modified pEX3 expression vector. The expression plasmid containing the

DNA from IgE constant region exon 3 was called ΔExce34 (the plasmid was also called deltaEXCH3, and the fusion protein produced was called the deltaEXCH3 protein).

Recombinant IgE plasmids AExce34 were introduced into 5 E. coli strain N4830-1 by transformation (this cell line was called AExce34/E. coli N4830, and is deposited with the ATCC as Accession No.: (not yet assigned)). growth of the bacteria overnight at 28°C, recombinant protein expression was induced by subculture 10 prewarmed medium at 42°C and incubation for 2 hours.

Purification of recombinant protein was accomplished using the following method. Induced cell pellets were sonicated successively on ice in the following solutions: TEN (0.1 M Tris-Cl, pH7.0, 10 mM EDTA, 0.15M NaCl);

TET (50 mM Tris-Cl, pH7.5, 0.5 mM EDTA, 2% Triton X-100); TED (50 mM Tris-Cl, pH7.5, 0.5 mM EDTA, 2% sodium deoxycholate); TU (50 mM Tris-Cl, pH7.5, 2 M urea). sonication in each solution, lysates were incubated 30 minutes on ice and insoluble recombinant proteins were sedimented by centrifugation at 10,000 x G for 10 minutes at 4°C. The final pellet was dissolved in TU9DTT (50mM Tris-Cl pH 7.5, 9M urea, 1 mM Dithiothreitol) sonication at room temperature; was incubated 30 minutes with shaking at room temperature; and, the dissolved 25 protein was clarified by centrifugation as above at room temperature. The supernatant was adjusted to 1 mg/ml and dialyzed against three changes of 100 volumes of 10 mM KPO. for three days. The purified protein was evaluated for

The

size of the

purity by gel electrophoresis.

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recombinant fusion protein was as predicted for a protein comprised of 64 amino acid residues of IgE C_R3 and the approximately 400 residues from β -galactosidase. Final proof that the protein contained canine IgE heavy chain constant region was determined based on cross-reactivity of monoclonal antibodies raised to authentic canine IgE with the recombinant protein.

One skilled in the art will appreciate that many different cloning vectors could be used to express canine IgE polypeptides, or to generate recombinant cell lines comprising canine IgE genomic material. The present example is an illustrative embodiment. The invention described herein comprises the cloning and cloning and expression of canine IgE heavy chain constant region material in various cloning and cloning-expression systems, including those derived from both prokaryotic or eukaryotic cells and organisms, those using other types of cloning vectors such as other plasmids, bacteriophage or viruses that replicate in eukaryotic cells.

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Examples

The DNA sequence for canine IgE heavy chain constant region, as described herein, has been used to develop methods to facilitate the treatment of canine allergies, including Type I immediate-hypersensitivity reactions, see, e.g., copending U.S. application Serial No. (not yet assigned), entitled Method and Compositions to Facilitate Treatment of Allergy in Dogs, filed 8 December 1995, in the name of MacKinnon et al, which is incorporated by

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reference herein. The invention is also used to treat canine allergies by production of recombinant peptides, so as to achieve canine immunomodulation. These peptides are 5 administered with art recognized pharmacologic excipients; these pharmacologic compositions are administered in pharmacologic dosages determined in accordance with skill known to those of ordinary skill in the art in view of the clinical presentation of the dog, to achieve canine immunomodulation.

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Example 1: An antisense or a sense sequence to a DNA sequence encoding canine IgE heavy chain constant region is used to screen the genomic material of a non-canine species. Such screening identifies the region of the 15 genome in the non-canine species that encodes IgE heavy chain constant region, where this IgE region is not yet In a preferred embodiment, the PCR walking approach of the present invention is employed to advantage in isolating IgE regions of a species.

20 Moreover, an antisense or a sense sequence to a canine IgE heavy chain constant region DNA sequence disclosed herein is used to determine the extent of homology between the canine IgE sequence and the sequence an isolated noncanine IgE gene. Knowledge 25 interspecies homology, in combination with biological data, is used to locate functional regions in the IgE molecule of either species.

For example, it is determined that canine IgE and feline IgE both bind to the feline IgE receptor, thus the nucleic acid residues encoding the amino acids responsible for this binding would likely be found among those which

are homologous between the canine and feline sequences. Knowledge of the specific DNA sequence encoding the amino acids responsible for such binding is used to develop therapeutic compounds that inhibit the binding. For example, this is done by expressing a peptide comprising a binding domain which will compete with the binding of IgE to mast cells and basophils.

Additionally, it is determined that canine IgE but not feline IgE, bind to the canine IgE receptor. again, knowledge of interspecies sequence comparison is 10 Pursuant to this knowledge of the extent of interspecies homology between dog and feline genomes, canine/feline IqE are developed. Various canine/feline IgE hybrids are analyzed to determine which bind to the canine receptor. The region of the canine sequence which encodes peptide involved in receptor binding is determined by comparison of the hybrids that do bind relative to those that do not bind to the canine receptor.

Example 2: Monoclonals were raised to a recombinant peptide comprising a component, preferably less than total, of canine IgE heavy chain constant region peptide. The monoclonals were screened to determine reactivity with native canine IgE. A monoclonal was found which was reactive with native canine IgE. Accordingly, this monoclonal was used to diagnose canine allergy in accordance with standard diagnostic techniques.

The two primary sources of nonspecificity present in the art by use of polyclonal serum to detect allergen-

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specific canine IgE: 1) lack of specificity in detecting only IgE, and 2) the possibility of detecting any glycosylated antibody, are overcome by use of the present invention. These sources of nonspecificity were overcome by using a nonglycosylated component of IgE as a recombinant immunogen in order to develop monoclonal antibodies. Nonspecificity due to reactivity with any glycosylated antibody is overcome since when recombinant canine IgE is expressed in bacteria, it is nonglycosylated because bacteria do not glycosylate proteins.

Example 3: Heretofore, it has been believed that expression of canine IgE DNA in eukaryotic cells would require that the entire IgE molecule including the entire heavy chain constant region, as well as the heavy chain variable regions and the light chains. Although there have been reports that functional non-canine IgE peptides have been expressed, these reports have generally been anecdotal and no consistently reproducible protocols have been reported. No known functional canine IgE peptide has ever been reported. There is only one known report of expression of canine IgE DNA, wherein it was expressed as part of a chimeric mouse antibody. See, e.g., Chang, U.S. Patent 5,514,776, issued 7 May 1996.

As disclosed herein for the first time in the art, it

25 has been found that canine IgE having less than the full
constitution of light and heavy chains can be expressed in
eukaryotic cells. Moreover, it has been found that
proteins expressed thereby possess functional qualities of
the canine IgE molecule. Accordingly, it is disclosed

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that the region on the canine IgE molecule that is bound by the canine IgE receptor is contained within the peptide sequence encoded by exons CH2 through CH4.

A strategy for the rational design of therapeutic agents based on recombinant proteins is to develop a variant of a naturally occurring protein which retains some but not all of the properties of the authentic protein. For canine IgE, two relevant functions are its binding to the IgE receptor and its association with allergen. A variant recombinant IgE which retains the binding functions but can not associate with allergen is of therapeutic value within this context. In order for IgE to associate with allergen, the IgE variable domains are needed.

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15 Based the sequences disclosed herein, recombinant protein was developed that lacked the ability to associate with allergen but possessed other native canine IqE characteristics. Accordingly, a recombinant protein derived from canine IgE regions CH2, CH3, and CH4 20 developed. Recombinant DNA was designed constructed so as to fuse a signal (secretion) sequence to canine IqE exons 2, 3, and 4. The signal secretion sequence is set forth in figure 5 together with canine IgE heavy chain sequences for exons 2-4. This recombinant DNA was inserted into a mammalian expression vector pcDNA3 25 (INVITROGEN, San Diego, CA) and the expression vector was cells mammalian (COS cells) introduced into by Supernatants from the transfected cells transfection. were evaluated by ELISA analysis and western blot. The

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results indicated that the transfected cells produced a recombinant IqE which was secreted as a dimeric molecule. This secreted IgE polypeptide had an amino acid sequence encoded by canine IgE exons 2-4; this polypeptide was found to bind to a recombinant canine IgE receptor in vitro. Thus, it was determined that the canine IqE receptor binds a portion of the canine IgE molecule encoded by exons 2-4. The portion of the canine IgE molecule bound by the IgE receptor is herein termed the 10 "Fc region." The truncated Fc region protein cannot associate with allergen because it lacks IqE variable Since this truncated recombinant IgE protein regions. cannot associate with allergen, it is used as an antiallergy therapeutic agent. A recombinant IgE Fc region 15 protein with identical properties was produced by insect cells infected with recombinant baculovirus, and is also used as an anti-allergy therapeutic agent.

Accordingly, a recombinant canine IgE molecule comprising protein as disclosed herein is produced that 20 comprises a region corresponding to the Fc region of canine IgE while lacking the allergen-binding components of the native IgE molecule. This peptide is administered in a pharmaceutical form to a dog experiencing allergic disease. The dosage administered is modulated accordance with skill held by one of ordinary skill in the art in view of the clinical manifestations of the patient. Such recombinant peptides bind to the surfaces of mast cells and basophils by the canine IgE Fc receptors on The binding of the recombinant peptides these cells.

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competes with and precludes the binding of circulating canine IgE to the surfaces of these cells. Since the recombinant peptides do not have allergen-binding regions, allergen cannot be bound, and anaphylatoxin release is diminished.

Example 4: As described in Example 3, a recombinant canine IgE nucleic acid sequence corresponding to less than the complete canine IgE heavy chain constant and variable regions together with the canine IgE light chains 10 was produced which led to a secreted recombinant IgE peptide fragment; this fragment retained several functions of the authentic IgE. Similarly, a recombinant IgE corresponding to: CH1, CH2, CH3, or CH4 combination thereof); and, CH5 and/or CH6 was produced by 15 mammalian cells transfected with the corresponding recombinant canine DNA. These cells are evaluated for the expression of cell-bound IgE. A cell-line expressing this truncated surface IqE is used to raise antibodies which bind to IgE on the surface of memory B-cells, monoclonal 20 or polyclonal antibodies are raised for this purpose.

Accordingly, glycosylated (e.q., expressed eukaryotic cells) recombinant canine IgE heavy chain epsilon protein is used to produce a canine antibody to immunomodulation; monoclonal orpolyclonal achieve 25 antibodies are used for this purpose. IgE-producing Blymphocytes (memory B cells) have IgE molecules on their surfaces. The surface IgE is membrane bound due to amino acid residues encoded by exons 5 and 6. The amino acid residues encoded by exons 5 or 6 are not part of secreted

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IgE heavy chain constant region which is completely encoded by exons 1, 2, 3 and 4. The amino acids encoded by exon 5 are not entirely encompassed by the membrane; a 5 component of the peptide encoded by this exon accessible in the extramembrane environment.

For example, a cell-line is developed that expresses recombinant IgE polypeptide comprising peptide encoded by exon 5 and/or exon 6. The polypeptide can optionally comprise peptide encoded by canine IgE heavy chain epsilon exons 1, 2, 3, 4, or 6. The membrane-bound peptide produced is used as an immunogen in the production, e.g., of monoclonal antibodies. The monoclonals raised to such immunogen are screened. Monoclonals that inhibit the 15 binding of membrane-bound IgE to a material to which the IgE has binding affinity are chosen; monoclonals are administered to achieve immunomodulation e.q. by lessening IqE production by memory B cells. Alternatively, monoclonals are chosen which are cytotoxic to memory B cells that express IgE. 20

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Antibodies are chosen which bind to membrane-bound IgE not other IgE molecules to any adverse degree. Adverse binding to IgE populations which are not membranebound may lead to an undesired decrease in an animal's defense to parasitic organisms.

These monoclonals are administered to doa experiencing allergic disease. These monoclonal antibodies treat allergic disease in dogs by decreasing the extent of Type I hypersensitivity reactions. hypersensitivity reactions are diminished by lessening the binding of allergens to membrane-bound IgE on memory B

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Absent immunomodulation of this type, allergens would bind to membrane-bound IgE and this binding leading to production of circulating IqE that would bind to the 5 surface of mast cells and basophils by Fc receptors on these cells, the Fc receptor-bound IqE would then bind to allergens leading to the release of anaphylaxis-inducing substances, such as vasoactive amines.

Example 5: A truncated recombinant IgE described in 10 Example 3 and Example 4 is used to develop antibodies which react with authentic canine IgE. Bacterial-derived recombinant canine IqE peptides are also Accordingly, a recombinant canine IgE heavy chain constant protein in accordance with the invention (glycosylated or nonglycosylated) is produced that comprises a peptide region corresponding to the Fc region Monoclonal antibodies are raised to this of canine IgE. A pharmaceutical is prepared which comprises such monoclonal antibodies and a pharmaceutical excipient. 20 The pharmaceutical is administered to a dog experiencing allergic disease in a dosage determined by one skilled in the art based on the clinical manifestations of the dog. The monoclonal antibody binds to the Fc region of the dog's circulating IgE and prevents binding of 25 circulating IgE to mast cells or basophils which would have lead to type I hypersensitivity reactions.

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Example 6: Scientific processes to develop variant peptides with improved (alternate) function include: By use of the sequence information disclosed herein

polypeptides are produced; one identifies peptide segments within said polypeptide which retain biological activity for some, but not all of the activity of the authentic polypeptide. In accordance with standard methodologies, 5 and based on the sequence of such peptide segments, peptide analog libraries are developed which contain alterations at various locations within the peptide Screening of these libraries identification of second generation peptides with improved or alternate function. (See, e.g., Gordon EM, et al., "Applications of Combinatorial Technologies to Drug Discovery I., " J. Med. Chem. 37:1233-1251 (1994).)

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The sequence of the active peptides is Example 7: utilized to design non-peptide analogs, often referred to as "small molecules." In accordance with methodologies known in the art, the chemical nature of the amino acid residues comprising the disclosed peptide or segments thereof is evaluated; this evaluation is used to determine organic compounds which contain analogous juxtapositioning of these chemical groups. (See, e.g., Smith, R.G., et al. Nonpeptidyl Growth Hormone Secretoque," Science 260:1640 (1993).)

Example 8: Polypeptides in accordance with those disclosed herein are used to prepare a model that mimics the binding of canine IgE to IgE receptor; upon the disclosure herein, the model is prepared in accordance with methodologies known to those of ordinary skill in the This model is used to screen pharmaceutical compounds. Pursuant to this screening, compounds are

selected that have an affect on the extent of binding of IgE to receptor.

Example 9: Antisense reagents are oligonucleotides that are capable of entering a cell and binding to the sense mRNA. When the antisense reagents bind to the target sense mRNA, they block expression of the peptide encoded by the sense sequence. A further application of the sequence information disclosed herein comprises use of "antisense" reagents that specifically target cells which 10 produce canine IqE messenger RNA. Antisense reagents based on the sequences of exons 1-4 target all producing cells, and impair production of any IgE whether secreted or bound. Antisense reagents based on the sequences of exons 5 and 6 specifically target the memory 15 B-cells that produce surface-bound IgE, and impair production of the bound IgE.

Closing

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It will be appreciated by one skilled in the art that in addition to the DNA sequence for canine IgE heavy chain constant region disclosed herein, conservative variants are also effective for various utilities; accordingly, the invention comprises nontotal nucleotide sequences in accordance with those disclosed herein, as well as nucleotide sequences that have less than total homology but do exhibit specific hybridization with a sequence or segment thereof disclosed herein. Conservative variants include nucleotide substitutions that do not result in changes in the amino acid sequence as well as nucleotide

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substitutions that result in conservative amino acid substitutions, or amino acid substitutions which do not substantially affect the character of the polypeptide translated from said nucleotides. For example, polypeptide character is not substantially affected if the substitutions do not preclude specific binding of the peptide to canine IgE receptor or other canine IgE ligands.

All publications mentioned herein are incorporated herein by reference to describe and disclose specific information for which the reference was thus discussed. It is to be noted that as used herein and in the appended claims, the singular forms "a" and "the" include plural referents unless the context clearly indicates otherwise. Thus, for example, reference to "a formulation" includes 15 mixtures of different formulations and reference to "the method of treatment" includes reference to equivalent steps and methods known to those skilled in the art, and Unless defined otherwise, all technical and scientific terms used herein have the same meaning as 20 commonly understood by one of ordinary skilled in the art. Although methods and materials or equivalent to those described herein can be used in the practice for testing of the invention, the preferred methods and materials are 25 described herein. It is understood that the invention is limited solely be the appended claims.

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Sequence Listing

- (1) GENERAL INFORMATION
- (i) APPLICANT: IDEXX Laboratories, Inc.
- 5 (ii) TITLE OF INVENTION: Recombinant DNA Molecules That Express
 Canine IgE Heavy Chain Constant Region and Components
 Thereof
 - (iii) NUMBER OF SEQUENCES: 2
 - (iv) CORRESPONDENCE ADDRESS:
- 10 (A) ADDRESSEE: Lyon & Lyon
 - (B) STREET: 633 West Fifth Street
 - (C) CITY: Los Angeles
 - (D) STATE: California
 - (E) COUNTRY: USA
- 15 (F) ZIP: 90071-20066
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- 20 (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- 25 (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Consalvi, Mary S.
 - (B) REGISTRATION NUMBER: 32,212
 - (C) REFERENCE/DOCKET NUMBER: 213/223
 - (ix) TELECOMMUNICATION INFORMATION:
- 30 (A) TELEPHONE: 213-489-1600
 - (B) TELEFAX: 213-955-0440

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	(2)	INFORMATION FOR SEQUENCE ID NO: 1:	
	(i)	SEQUENCE CHARACTERISTICS	
		(A) LENGTH: 1842 base pairs	
5		(B) TYPE: nucleic acid	
		(C) STRANDEDNESS: double	
		(D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
10			
	GTCC	AGTGAC CTCCATCTCT GCCCCCATGC TTTTCCTTCT	40
	CAGA	CGCCCC CTGGGGCCAG GAGCAGGATA CCCCAGGTCA	80
	ACAG	CGGGCC TGGCATATGA TGGGGTGACA GTCCCAAGGC	120
	AGGC	ACTGAC ACTGGNCCTG TCCCCACAGC CACCAGCCAG	160
15	GACCT	rg	166
		STG TTC CCC TTG GCC TCC TGC TGT AAA GAC	199
	Ser V	Val Phe Pro Leu Ala Ser Cys Cys Lys Asp	
20	AAC A	ATC GCC AGT ACC TCT GTT ACA CTG GGC TGT	232
		Ile Ala Ser Thr Ser Val Thr Leu Gly Cys	
		are the ser and ser the mod cry cys	
	CTG (STC ACC GGC TAT CTC CCC ATG TCG ACA ACT	265
	Leu V	Val Thr Gly Tyr Leu Pro Met Ser Thr Thr	
25			
	GTG A	ACC TGG GAC ACG GGG TCT CTA AAT AAG AAT	298
	Val 7	Thr Trp Asp Thr Gly Ser Leu Asn Lys Asn	
	GTC A	ACG ACC TTC CCC ACC ACC TTC CAC GAG ACC	331
30	Val 1	Thr Thr Phe Pro Thr Thr Phe His Glu Tyr	

TAC GGC CTC CAC AGC ATC GTC AGC CAG GTG ACC

	Thr Gly Leu His Ser Ile Val Ser Gln Val Thr	
	GCC TCG GGC GAG TGG GCC AAA CAG AGG TTC ACC	397
	Asp Ser Gly Glu Trp Ala Lys Gln Arg Phe Thr	
5		
	TGC AGC GTG GCT CAC NNT GAG TCC ACC GCC ATC	430
	Cys Ser Val Ala His Xaa Glu Ser Thr Ala Ile	
10	AAC AAG ACC TTC AGT GGT	448
	Asn Lys Thr Phe Ser Ala	
		400
	AANCCAGGGT TNNNTGGCCA CATGACACTG GAGGGAGAAG GGACAGGCTG GNGAATGCGC CATGGCTGGT AACGCCCAGC	488 528
15	ANATGTGGGG CTGGGGCTGA CACATGAGTC CCGTGGGCTN	568
	AGAGACACCA CTGCCACATG GCTGCCTCTA CTTCTAGCA	607
	TGT GCC TTA AAC TTC ATT CCG CCT ACC GTG AAG	640
	Cys Ala Leu Asn Phe Ile Pro Pro Thr Val Lys	
20		
	CTC TTC CAC TCC TCC TGC AAC CCC GTC GGT GAT	673
	Leu Phe His Ser Ser Cys Asn Pro Val Gly Asp	
	ACC CAC ACC ACC ATC CAG CTC CTG TGC CTC ATC	706
25	Thr His Thr Thr Ile Gln Leu Leu Cys Leu Ile	700
	0	
	TCT GGC TAC GTC CCA GGT GAC ATG GAG GTC ATC	739
	Ser Gly Tyr Val Pro Gly Asp Met Glu Val Ile	
30	TGG CTG GTG GAT GGG CAA AAG GCT ACA AAC ATA	772
	Trp Leu Val Asp Gly Gln Lys Ala Thr Asn Ile	

	TTC	CCA	TAC	ACT	GCA	CCC	GGC	ACA	AAG	GAG	GGC		805
	Phe	Pro	Tyr	Thr	Ala	Pro	Gly	Thr	Lys	Glu	Gly		
	AAC	GTG	ACC	TCT	ACC	CAC	AGC	GAG	CTC	AAC	ATC		838
5	Asn	Val	Thr	Ser	Thr	His	Ser	Glu	Leu	Asn	Ile		
	ACC	CAG	GGN	NNG	TGN	GTA	TCC	CAA	AAA	ACC	TAC		871
	Thr	Gln	Gly	Xaa	Trp	Val	Ser	Gln	Lys	Thr	Tyr		
10	ACC	TGC	CAG	GTC	ACC	TAT	CAA	GGC	TTT	ACC	TTT		904
	Thr	Cys	Gln	Val	Thr	Tyr	Gln	Gly	Phe	Thr	Phe		
	AAA	GAT	GAG	GCT	CGC	AAG	TGC	TCA	GGT				931
15	Lys	Asp	Glu	Ala	Arg	ГÀЗ	Суз	Ser	Glu				
	ATG	3CCC(CCC 1	rgtco	CCCI	AG AA	AACC	CAGAT	C GCC	CGAC	GCT		971
	CAG	AGATO	AG (GCC2	AAGG	CA CO	GCC:	CATO	CAC	CCT	CTCA	1	011
	CAC	ACTG	CAG A	AG								1	.023
20	TCC	GAC	CCC	CGA	GGC	GTG	AGC	AGC	TAC	CTG	AGC	1	.056
	Asp	Tyr	Pro	Arg	Gly	Val	Ser	Ser	Tyr	Leu	Ser		
	CCA	CCC	AGC	CCC	CTT	GAC	CTG	TAT	GTC	CAC	AAG	1	.089
	Pro	Pro	Ser	Pro	Leu	Asp	Leu	Tyr	Val	His	Lys		
25	GCG	CCC	AAG	ATC	ACC	TGC	CTG	GTA	GTG	GAC	CTG	1	.122
	Ala	Pro	Lys	Ile	Thr	Сув	Leu	Val	Val	Asp	Leu		
	GCC	ACC	ATG	GAA	GGC	ATG	AAC	CTG	ACC	TGG	TAC	1	155
30	Ala	Thr	Met	Glu	Gly	Met	Asn	Leu	Thr	Trp	Tyr		
	CGC	GAG	AGC	ΔΔΔ	GAA	CCC	GTG	אאר	CCG	GTC	ССТ	1	188

	Arg Glu Ser Lys Glu Pro Val Asn Pro Val Pro	
	TTG AAC AAG AAG GAT CAC TTC AAT GGG ACG ATC	122
	Leu Asn Lys Lys Asp His Phe Asn Gly Thr Ile	
5		
	ACA GTC ACG TCT ACC CTG CCA GTG AAC ACC AAT	1254
	Thr Val Thr Ser Thr Leu Pro Val Asn Thr Asn	
	GAC TGG ATC GAG GGC GAG ACC TAC TAT TGC AGG	
10		1287
10	Asp Trp Ile Glu Gly Glu Thr Tyr Tyr Cys Arg	
	GTG ACC CAC CCG CAC CTG CCC AAG GAC ATC GTG	1320
	Val Thr His Pro His Leu Pro Lys Asp Ile Val	
15		
	CGC TCC ATT GCC AAG GCC CCT GGT	1344
	Arg Ser Ile Ala Lys Ala Pro Gly	
	·	
0.0	GAGCCACGGG CCCAGGGGAG GTGGGCGGGC CTCCTGANCC	1384
20	GGAGCCTGGG CTGACCCCAC ACCTATCCAC AGGC	1418
	AAG CGT GCC CCC CCG GAT GTG TAC TTG TTC CTG	
	Lys Arg Ala Pro Pro Asp Val Tyr Leu Phe Leu	1451
	1 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -	
25	CCA CCG GAG GAG GAG CAG GGG ACC AAG GAC AGA	1484
	Pro Pro Glu Glu Glu Gln Gly Thr Lys Asp Arg	
	GTC ACC CTC ACG TGC CTG ATC CAG AAC TTC TTC	1517
	Val Thr Leu Thr Cys Leu Ile Gln Asn Phe Phe	
30		
	CCC GAG GAC ATT TCA GTG CAA TGG CTG CGA AAC	1550
	Pro Glu Asp Ile Ser Val Gln Trp Leu Arg Asn	

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	GAC AGC CCC ATC CAG ACA GAC CAG TAC ACC ACC	1583
	Asp Ser Pro Ile Gln Thr Asp Gln Tyr Thr Tyr	
	ACG GGG CCC CAC AAG GTC TCG GGC TCC AGG CCT	1616
5	Thr Gly Pro His Lys Val Ser Gly Ser Arg Pro	
	GCC TTC TTC ATC TTC AGT CGC CTG GTG GAC TGG	1649
	Ala Phe Phe Ile Phe Ser Arg Leu Val Asp Trp	
10	GAG CAG AAA AAC AAA TTC ACC TGC CAA GTG GTG	1682
	Glu Gln Lys Asn Lys Phe Thr Cys Gln Val Val	
	CAT GAG GCG CTG TCC GGC TCT AGG ATC CTC CAG	1715
15	His Glu Ala Leu Ser Gly Ser Arg Ile Leu Gln	
15		
	AAA TGG GTG TCC AAA ACC CCC GGT AAA	1742
	Lys Trp Val Ser Lys Thr Pro Gly Lys	
20	TGATGCCCAC CCTCCTCCCG CCGCCACCCC CCAGGGCTCC	1782
	ACCTGCTGGG GCAGGGGAGG GGGGCTGGCA AGACCCTCCA	1822
	TCTATCCTTN TCAATAAACA	1842
	(2) INFORMATION FOR SEQUENCE ID NO: 2:	
25		
	(i) SEQUENCE CHARACTERISTICS	
	(A) LENGTH: 538 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: double	
30	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

	GGGGAGGGG GGCGGGTCTG CCTTCCCCCN ACCAGCACAT	40
	GAACGGCTGG ACCGGGGAGG GNTGACTGGC CGGTGCCCGC	80
	A	81
5	GAG CTA GAG CTC CAG GAG CTG TGC GCG GAT GCC	114
	Glu Leu Glu Leu Gln Glu Leu Cys Ala Asp Ala	
	NOTE CALL ACTUAL CALL COME CALL CALL	
	ACT GAG AGT GAG GAG CTG GAC GAG CTG TGG GCC	147
10	Thr Glu Ser Glu Glu Leu Asp Glu Leu Trp Ala	
10	AGC CTG CTC ATC TTC ATC ACC CTC TTC CTG CTC	100
	Ser Leu Leu Ile Phe Ile Thr Leu Phe Leu Leu	180
	and the sea the sea the sea sea	
	AGA GTG AGC TAC GGC GCC ACC AGC ACC CTC TTC	213
15	Arg Val Ser Tyr Gly Ala Thr Ser Thr Leu Phe	
	AAG	216
	Lys	
20		
	GTGGGCATGC AGAGCCCCTG GCCGGGGGTG GGGGCAGCAC	256
	AGAGGGAGNG AGAGGTCCCG GCAGAGCTGT CCTCACATGT	296
	GCCCTCCCCC CAGGTGAAG	315
25	TGG GTA CTC GCC ACC GTC CTG CAG GTG AAG CCA	240
	Trp Val Leu Ala Thr Val Leu Gln Val Lys Pro	348
	The same and the det can var by the	
	CAG GCC GCC CAA GAC TAC GCC AAC ATC GTG CGG	381
	Gln Ala Ala Gln Asp Tyr Ala Asn Ile Val Arg	
30		
	CCG GCA CAG	390
	Pro Ala Cln	

TAGGCCCAGA	GACACGGTGA	CGAGGCCTTG	CTTTCTGCCC	430
CCCNNNNNCC	GGCTGAGGGC	AATCTGCTGG	CCCTGAGTGG	470
GAGGAGGAAA	GCAGACAAAC	NCAGAGGGGC	CAGAGCCAGA	510
CGCCCAGCAC	ACACGGATCC	AGAAGCTT		538

Claims

- An isolated DNA sequence comprising: a DNA 1. sequence of canine IgE heavy chain constant region exon 1 (SEQ ID NO: 1: nucleotides 167-448), or conservative 5 variants thereof; a DNA sequence of canine IgE heavy chain constant region exon 2 (SEQ ID NO: 1: nucleotides 608-931), or conservative variants thereof; a DNA sequence of canine IgE heavy chain constant region exon 3 (SEQ ID NO: nucleotides 1024-1344), or conservative variants 10 thereof; a DNA sequence of canine IgE heavy chain constant region exon 4 (SEQ ID NO: 1: nucleotides 1419-1742), or conservative variants thereof; a DNA sequence of canine IgE heavy chain constant region exon 5 (SEQ ID NO: 2: nucleotides 82-216), or conservative variants thereof; or, 15 a DNA sequence of canine IgE heavy chain constant region NO: 2: nucleotides 316-390), (SEO ID conservative variants thereof.
- A purified polypeptide comprising an amino acid
 sequence encoded by a nucleic acid sequence in accordance with claim 1.
 - 3. A conservative variant of a polypeptide in accordance with claim 2.
 - 4. A recombinant polypeptide of claim 2 produced by a eukaryotic cell.

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5. A segment of a nucleotide sequence in accordance with claim 1 comprising eighteen contiguous nucleotides in a reading frame of: a DNA sequence of canine IgE heavy chain constant region exon 1 (SEQ ID NO: 1: nucleotides 167-448); a DNA sequence of canine IgE heavy chain constant region exon 2 (SEQ ID NO: 1: nucleotides 608-931); a DNA sequence of canine IgE heavy chain constant region exon 3 (SEQ ID NO: 1: nucleotides 1024-1344); a DNA sequence of canine IgE heavy chain constant region exon 4 (SEQ ID NO: 1: nucleotides 1419-1742); a DNA sequence of canine IgE heavy chain constant region exon 5 (SEQ ID NO: 2: nucleotides 82-216); or, a DNA sequence of canine IgE heavy chain constant region exon 6 (SEQ ID NO: 2: nucleotides 316-390).

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- 6. A nucleotide sequence complementary to a sequence of claim 5.
- 7. A segment of a nucleotide sequence in accordance 20 with claim 5 comprising twenty-one contiguous nucleotides.
 - 8. A prokaryotic or eukaryotic cloning vector comprising a DNA sequence according to claim 1.
- 9. A prokaryotic vector of claim 8 wherein said vector is pBKce5 or Δexce34.
 - 10. A eukaryotic vector of claim 8 wherein said vector expresses a polypeptide having a sequence encoded

by less than canine IgE heavy chain constant region exon 1 (SEQ ID NO: 1: nucleotides 167-448), or conservative variants thereof; the DNA sequence of canine IqE heavy chain constant region exon 2 (SEQ ID NO: 1: nucleotides 5 608-931), or conservative variants thereof; the DNA sequence of canine IgE heavy chain constant region exon 3 (SEQ ID NO: 1: nucleotides 1024-1344), or conservative variants thereof; the DNA sequence of canine IqE heavy chain constant region exon 4 (SEQ ID NO: 1: nucleotides 10 1419-1742), or conservative variants thereof; the DNA sequence of canine IgE heavy chain constant region exon 5 (SEQ ID NO: 2: nucleotides 82-216), or conservative variants thereof; and, the DNA sequence of canine IgE heavy chain constant region exon 6 (SEQ ID NO: 2: 15 nucleotides 316-390), or conservative variants thereof.

- 11. A polypeptide encoded by a vector of claim 8.
- 12. A polypeptide encoded by a eukaryotic vector of 20 claim 8.
 - 13. A pharmaceutical composition comprising a polypeptide encoded by a vector of claim 8 and a pharmaceutical excipient.

- 14. An antibody raised to a polypeptide of claim 11.
- 15. An antibody of claim 14 for use in a method of treating the manifestations of allergy in a dog.

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- 16. A pharmaceutical composition comprising an antibody of claim 14 and a pharmaceutical excipient.
- 17. A eukaryotic vector of claim 8 wherein said vector expresses a polypeptide encoded by the DNA sequence of canine IgE heavy chain constant region exon 2 (SEQ ID NO: 1: nucleotides 608-931), or conservative variants thereof; the DNA sequence of canine IgE heavy chain constant region exon 3 (SEQ ID NO: 1: nucleotides 1024-1344), or conservative variants thereof; and, the DNA sequence of canine IgE heavy chain constant region exon 4 (SEQ ID NO: 1: nucleotides 1419-1742), or conservative variants thereof.
- 18. A sequence of claim 1 which comprises a nucleic acid sequence which encodes the Fc region of the canine IgE molecule or conservative variants thereof.
- 19. The sequence of claim 18 which consists 20 essentially of the DNA sequence of canine IgE heavy chain constant region exon 2 (SEQ ID NO: 1: nucleotides 608-931), or conservative variants thereof; the DNA sequence of canine IgE heavy chain constant region exon 3 (SEQ ID NO: 1: nucleotides 1024-1344), or conservative variants thereof; and, the DNA sequence of canine IgE heavy chain constant region exon 4 (SEQ ID NO: 1: nucleotides 1419-1742), or conservative variants thereof.
 - 20. A polypeptide encoded by a sequence of claim 18.

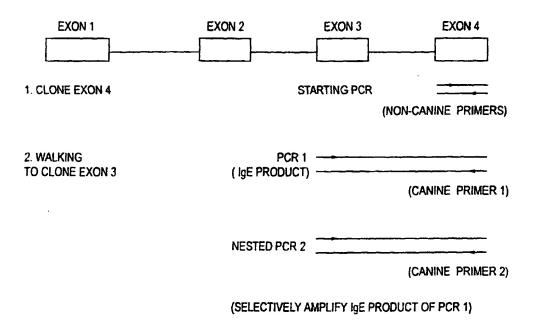
WO 97/30156

- 21. A polypeptide of claim 20 for use in a method of treating allergic manifestations in a dog.
- 22. A recombinant cell line comprising a DNA 5 sequence according to claim 1.
 - 23. A recombinant eukaryotic cell line in accordance with claim 22.
- 24. A recombinant cell line in accordance with claim22 wherein the cell line is ΔΕxce34/E. coli 4830.
- 25. A method for producing a polypeptide having an amino acid sequence homologous to a component of canine 15 IgE heavy chain constant region, said method comprising a step of expressing a peptide encoded by a cloning vector according to claim 8.
- 26. A cell that expresses a recombinant polypeptide 20 encoded by a DNA sequence of claim 1.
- 27. An antisense reagent capable of binding to sense mRNA encoded by a nucleic acid sequence of claim 1; whereby upon binding of the antisense reagent to the sense mRNA, production of the peptide encoded by the mRNA is impaired.
 - 28. The antisense reagent of claim 1 wherein the reagent impairs production of canine IgE.

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29. The antisense reagent of claim 28 wherein the reagent impairs production of membrane-bound canine IgE.

Fig. 1: NESTED WALKING PROCEDURE FOR CLONING CANINE IGE



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		HHCO1J Heavy constant chain (IGE CL') - Human MHCO2X Heavy constant chain (IGE a CL) - Mouse
		Translated sequence IDEXX IgE clone exons 1-4, spliced mRNA
Seq	1	SVFPLTRCCKNIPSNATSVTLGCLATGYFPEPVMVTWDTGSLNGTTMTLPATTLTLSGHY :: :: :: :: :: :: :: :: :: :: :: :: ::
Seq	2	PLKP CKGTA SMTLGCLVKDYFPGPVTVTWYSDSLNMSTVNFPALGSELK
Seq	3	SVFPLASCCKD-NIASTSVTLGCLVTGYLPMSTTVTWDTGSLNKNVTTFPTTEHFTYGLH
Seq	1	ATISLLTVSGAWAKQMFTCRVAHTPSSTDWVDNKTFSFTPPTVKILQSSCDGGGHFPP
Seq	2	VTTSQVTSWGKSAKN-FTCHVTHPPSFNESRTILVRPVNITEPTLELLHSSCDPNA-FHS
Seq	3	SIVSQVTASGEWAKQRFTCSVAHXESTAINKTFSACALNFIPPTVKLFHSSCNPVGDTHT
Seq	1	TIQLLCLVSGYTPGTINITWLEDGQVMDVDLS-TASTTQEGELASTQSELTLSQKHWLSD
Seq	2	TIQLYCFIYGHILNDVSVSWLMDDREITDTLAQTVLIKEEGKLASTCSKLNITEQQWMSE
Seq	3	TIQLLCLISGYVPGDMEVIWLVDGQKATNIFPYTAPGTKEGNVTSTHSELNITQXXXVSQ
Seq	1	RTYTCQVTYQGHTFEDSTKKCADSNPRGVSAYLSRPSPFDLFIRKSPTITCLVVDLAPSK
Seq	2	STFTCKVTSQGVDYLAHTRRCPDHEPRGVITYLIPPSPLDLYQNGAPKLTCLVVDLESEK
Seq	3	KTYTCQVTYQGFTFKDEARKCSESDPRGVSSYLSPPSPLDLYVHKAPKITCLVVDLATME
Seq	1	GTVNLTWSRASGKPVNHSTRKEEKQRNGTLTVTSTLPVGTRDWIEGETYQCRVTHPHLPR
Seq	2	N-VNVTWNQEKKTSVSASQWYTKHHNNATTSITSILPVVAKDWIEGYGYQCIVDHPDFPK
Seq	3	G-MNLTWYRESKEPVNPVPLNKKDHFNGTITVTSTLPVNTNDWIEGETYYCRVTHPHLPK
Seq	1	ALMRSTTKTSGPVGPRAAPEVYAFATP EWPGSRDKRTLACLIQNFMPEDISVQWLHNEV
Seq	2	PIVRSITKTPGQRSAPEVYVFPPPEEE SEDKRTLTCLIQNFFPEDISVQWLGDGK
Seq	3	DIVRSIAKAPGKRAPPDVYLFLPPEEEQGTKDRVTLTCLIQNFFPEDISVQWLRNDS
Seq	1	exon 4 * QLPDARHSTTQPRKTKGSGFFVFSRLEVTRAEWEQKDEFICRAVHEAASPSQTVQRAVS
Seq	2	LISNSQHSTTTPLKSNGSNQGFFIFSRLEVAKTLWTQRKQFTCQVIHEALQKPRKLEKTIS
Seq	3	PIQTDQYTTTGPHKVSGSRPAFFIFSRLVDWEQKNKFTCQVVHEALSGSRILQKWVS
Seq	1	VNPGK

Seq 2 TSLGN

Seq 3 KTPGK

Seq 1	1-420	Seq 1	1-420
Seq 2	1-407	Seq 3	1-417
Aligned	422	Aligned	426
Matches	185	Matches	228
Mismatches	237	Mismatches	198
Homology	43%	Homology	53%

FIG. 3A. CANINE IGE HEAVY CHAIN CONSTANT CHAIN DNA SEQUENCE WITH TRANSLATED EXONS 1 THRU 4 GTCCAGTGAC CTCCATCTCT GCCCCCATGC TTTTCCTTCT 40 CAGACGCCCC CTGGGGCCAG GAGCAGGATA CCCCAGGTCA 80 ACAGCGGCC TGGCATATGA TGGGGTGACA GTCCCAAGGC 120 AGGCACTGAC ACTGGNCCTG TCCCCACAGC CACCAGCCAG 160 GACCTG 166 TCT GTG TTC CCC TTG GCC TCC TGC TGT AAA GAC 199 Ser Val Phe Pro Leu Ala Ser Cys Cys Lys Asp AAC ATC GCC AGT ACC TCT GTT ACA CTG GGC TGT 232 Asn Ile Ala Ser Thr Ser Val Thr Leu Gly Cys CTG GTC ACC GGC TAT CTC CCC ATG TCG ACA ACT 265 Leu Val Thr Gly Tyr Leu Pro Met Ser Thr Thr GTG ACC TGG GAC ACG GGG TCT CTA AAT AAG AAT 298 Val Thr Trp Asp Thr Gly Ser Leu Asn Lys Asn GTC ACG ACC TTC CCC ACC ACC TTC CAC GAG ACC 331 Val Thr Thr Phe Pro Thr Thr Phe His Glu Tyr TAC GGC CTC CAC AGC ATC GTC AGC CAG GTG ACC 364 Thr Gly Leu His Ser Ile Val Ser Gln Val Thr GCC TCG GGC GAG TGG GCC AAA CAG AGG TTC ACC 397 Asp Ser Gly Glu Trp Ala Lys Gln Arg Phe Thr TGC AGC GTG GCT CAC NNT GAG TCC ACC GCC ATC 430 Cys Ser Val Ala His Xaa Glu Ser Thr Ala Ile AAC AAG ACC TTC AGT GGT 448 Asn Lys Thr Phe Ser Ala AANCCAGGGT TNNNTGGCCA CATGACACTG GAGGGAGAAG 488 GGACAGGCTG GNGAATGCGC CATGGCTGGT AACGCCCAGC 528 ANATGTGGGG CTGGGGCTGA CACATGAGTC CCGTGGGCTN 568 AGAGACACCA CTGCCACATG GCTGCCTCTA CTTCTAGCA 607 TGT GCC TTA AAC TTC ATT CCG CCT ACC GTG AAG 640 Cys Ala Leu Asn Phe Ile Pro Pro Thr Val Lys CTC TTC CAC TCC TCC TGC AAC CCC GTC GGT GAT 673 Leu Phe His Ser Ser Cys Asn Pro Val Gly Asp

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5/10 ACC CAC ACC ACC ATC CAG CTC CTG TGC CTC ATC Thr His Thr Thr Ile Gln Leu Leu Cys Leu Ile	706
TCT GGC TAC GTC CCA GGT GAC ATG GAG GTC ATC Ser Gly Tyr Val Pro Gly Asp Met Glu Val Ile	739
TGG CTG GTG GAT GGG CAA AAG GCT ACA AAC ATA Trp Leu Val Asp Gly Gln Lys Ala Thr Asn Ile	772
TTC CCA TAC ACT GCA CCC GGC ACA AAG GAG GGC Phe Pro Tyr Thr Ala Pro Gly Thr Lys Glu Gly	805
AAC GTG ACC TCT ACC CAC AGC GAG CTC AAC ATC Asn Val Thr Ser Thr His Ser Glu Leu Asn Ile	838
ACC CAG GGN NNG TGN GTA TCC CAA AAA ACC TAC Thr Gln Gly Xaa Trp Val Ser Gln Lys Thr Tyr	871
ACC TGC CAG GTC ACC TAT CAA GGC TTT ACC TTT Thr Cys Gln Val Thr Tyr Gln Gly Phe Thr Phe	904
AAA GAT GAG GCT CGC AAG TGC TCA GGT Lys Asp Glu Ala Arg Lys Cys Ser Glu	931
ATGGCCCCCC TGTCCCCCAG AAACCCAGAT GCGCGAGGCT CAGAGATGAG GGCCAAGGCA CGCCCTCATG CAGCCTCTCA CACACTGCAG AG	971 1011 1023
TCC GAC CCC CGA GGC GTG AGC AGC TAC CTG AGC Asp Tyr Pro Arg Gly Val Ser Ser Tyr Leu Ser	1056
CCA CCC AGC CCC CTT GAC CTG TAT GTC CAC AAG Pro Pro Ser Pro Leu Asp Leu Tyr Val His Lys	1089
GCG CCC AAG ATC ACC TGC CTG GTA GTG GAC CTG Ala Pro Lys Ile Thr Cys Leu Val Val Asp Leu	1122
GCC ACC ATG GAA GGC ATG AAC CTG ACC TGG TAC Ala Thr Met Glu Gly Met Asn Leu Thr Trp Tyr	1155
CGG GAG AGC AAA GAA CCC GTG AAC CCG GTC CCT Arg Glu Ser Lys Glu Pro Val Asn Pro Val Pro	1188
TTG AAC AAG AAG GAT CAC TTC AAT GGG ACG ATC Leu Asn Lys Lys Asp His Phe Asn Gly Thr Ile	1221

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ACA GTC ACG TCT ACC CTG CCA GTG AAC ACC AAT Thr Val Thr Ser Thr Leu Pro Val Asn Thr Asn	1254
GAC TGG ATC GAG GGC GAG ACC TAC TAT TGC AGG Asp Trp Ile Glu Gly Glu Thr Tyr Tyr Cys Arg	1287
GTG ACC CAC CCG CAC CTG CCC AAG GAC ATC GTG Val Thr His Pro His Leu Pro Lys Asp Ile Val	1320
CGC TCC ATT GCC AAG GCC CCT GGT	1344
Arg Ser Ile Ala Lys Ala Pro Gly	
GAGCCACGGG CCCAGGGGAG GTGGGCGGGC CTCCTGANCC GGAGCCTGGG CTGACCCCAC ACCTATCCAC AGGC	1384 1418
AAG CGT GCC CCC CCG GAT GTG TAC TTG TTC CTG Lys Arg Ala Pro Pro Asp Val Tyr Leu Phe Leu	1451
CCA CCG GAG GAG GAG CAG GGG ACC AAG GAC AGA Pro Pro Glu Glu Glu Gln Gly Thr Lys Asp Arg	1484
GTC ACC CTC ACG TGC CTG ATC CAG AAC TTC TTC Val Thr Leu Thr Cys Leu Ile Gln Asn Phe Phe	1517
CCC GAG GAC ATT TCA GTG CAA TGG CTG CGA AAC Pro Glu Asp Ile Ser Val Gln Trp Leu Arg Asn	1550
GAC AGC CCC ATC CAG ACA GAC CAG TAC ACC ACC ASp Ser Pro Ile Gln Thr Asp Gln Tyr Thr Tyr	1583
ACG GGG CCC CAC AAG GTC TCG GGC TCC AGG CCT Thr Gly Pro His Lys Val Ser Gly Ser Arg Pro	1616
GCC TTC TTC ATC TTC AGT CGC CTG GTG GAC TGG	1649
Ala Phe Phe Ile Phe Ser Arg Leu Val Asp Trp	
GAG CAG AAA AAC AAA TTC ACC TGC CAA GTG GTG Glu Gln Lys Asn Lys Phe Thr Cys Gln Val Val	1682
CAT GAG GCG CTG TCC GGC TCT AGG ATC CTC CAG His Glu Ala Leu Ser Gly Ser Arg Ile Leu Gln	1715
AAA TGG GTG TCC AAA ACC CCC GGT AAA Lys Trp Val Ser Lys Thr Pro Gly Lys	1742
TGATGCCCAC CCTCCTCCCG CCGCCACCCC CCAGGGCTCC ACCTGCTGGG GCAGGGAGG GGGGCTGGCA AGACCCTCCA TCTATCCTTN TCAATAAACA	1782 1822 1842

FIGURE 4: <u>Canine IgE heavy chain constant region DNA sequence v</u> <u>translated exons 5 and 6</u>	<u>with</u>
GGGGAGGGG GGCGGGTCTG CCTTCCCCCN ACCAGCACAT GAACGGCTGG ACCGGGGAGG GNTGACTGGC CGGTGCCCGC A	40 80 81
GAG CTA GAG CTC CAG GAG CTG TGC GCG GAT GCC Glu Leu Glu Leu Gln Glu Leu Cys Ala Asp Ala	114
ACT GAG AGT GAG GAG CTG GAC GAG CTG TGG GCC Thr Glu Ser Glu Glu Leu Asp Glu Leu Trp Ala	147
AGC CTG CTC ATC TTC ACC CTC TTC CTG CTC Ser Leu Leu Ile Phe Ile Thr Leu Phe Leu Leu	180
AGA GTG AGC TAC GGC GCC ACC AGC ACC CTC TTC Arg Val Ser Tyr Gly Ala Thr Ser Thr Leu Phe	213
AAG Lys	216
GTGGGCATGC AGAGCCCCTG GCCGGGGGTG GGGGCAGCAC AGAGGGAGNG AGAGGTCCCG GCAGAGCTGT CCTCACATGT GCCCTCCCCC CAGGTGAAG	256 296 315
TGG GTA CTC GCC ACC GTC CTG CAG GTG AAG CCA Trp Val Leu Ala Thr Val Leu Gln Val Lys Pro	348
CAG GCC GCC CAA GAC TAC GCC AAC ATC GTG CGG Gln Ala Ala Gln Asp Tyr Ala Asn Ile Val Arg	381
CCG GCA CAG Pro Ala Gln	390
TAGGCCCAGA GACACGGTGA CGAGGCCTTG CTTTCTGCCC CCCNNNNNCC GGCTGAGGGC AATCTGCTGG CCCTGAGTGG GAGGAGGAAA GCAGACAAAC NCAGAGGGGC CAGAGCCAGA CGCCCAGCAC ACACGGATCC AGAAGCTT	430 470 510 538

FIG. 5A Signal (secretion) sequence

M P A S M G G P A L L W

ECORI

GAATTCGGCC GCGAGATGCC TGCTTCGATG GGAGGCCCTG CCCTGCTGTG CTTAAGCCGG CGCTCTACGG ACGAAGCTAC CCTCCGGGAC GGGACGACAC

signal sequence cleavage

+1 L A L L L S S P G V M S S I A R A
5451 GCTAGCGCTG CTGCTCTCCT CTCCAGGTGT CATGTCATCA GCTCGTGCCT
CGATCGCGAC GACGAGAGGA GAGGTCCACA GTACAGTAGT CGAGCACGGA

start exon 2

- +1 L N F I P P T V K L F H S S C N P
 5501 TAAACTTCAT TCCGCCTACC GTGAAGCTCT TCCACTCCTC CTGCAACCCC
 ATTTGAAGTA AGGCGGATGG CACTTCGAGA AGGTGAGGAG GACGTTGGGG
- +1 V G D T H T T I Q L L C L I S G Y
 5551 GTCGGTGATA CCCACACCAC CATCCAGCTC CTGTGCCTCA TCTCTCGCTA
 CAGCCACTAT GGGTGTGGTG GTAGGTCGAG GACACGGAGT AGAGACCGAT
- +1 V P G D M E V I W L V D G Q K A
 5601 CGTCCCAGGT GACATGGAGG TCATCTGGCT GGTGGATGGG CAAAAGGCTA
 GCAGGGTCCA CTGTACCTCC AGTAGACCGA CCACCTACCC GTTTTCCGAT
- +1 T N I F P Y T A P G T K E G N V T
 5651 CAAACATATT CCCATACACT GCACCCGGCA CAAAGGAGGG CAACGTGACC
 GTTTGTATAA GGGTATGTGA CGTGGGCCGT GTTTCCTCCC GTTGCACTGG
- +1 S T H S E L N I T Q G X W V S Q K 5701 TCTACCCACA GCGAGCTCAA CATCACCCAG GGNNNGTGNG TATCCCAAAA AGATGGGTGT CGCTCGAGTT GTAGTGGGTC CCNNNCACNC ATAGGGTTTT
- +1 T Y T C Q V T Y Q G F T F K D E 5751 AACCTACACC TGCCAGGTCA CCTATCAAGG CTTTACCTTT AAAGATGAGG TTGGATGTGG ACGGTCCAGT GGATAGTTCC GAAATGGAAA TTTCTACTCC
- +1 A R K C S E S D P R G V S S Y L S 5801 CTCGCAAGTG CTCAGAGTCC GACCCCCGAG GCGTGAGCAG CTACCTGAGC GAGCGTTCAC GAGTCTCAGG CTGGGGGCTC CGCACTCGTC GATGGACTCG
- +1 P P S P L D L Y V H K A P K I T C
 5851 CCACCCAGCC CCCTTGACCT GTATGTCCAC AAGGCGCCCA AGATCACCTG
 GGTGGGTCGG GGGAACTGGA CATACAGGTG TTCCGCGGGT TCTAGTGGAC

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- +1 L V V D L A T M E G M N L T W Y
 5901 CCTGGTAGTG GACCTGGCCA CCATGGAAGG CATGAACCTG ACCTGGTACC
 GGACCATCAC CTGGACCGGT GGTACCTTCC GTACTTGGAC TGGACCATGG
- +1 R E S K E P V N P V P L N K K D H
 5951 GGGAGACCAA AGAACCCGTG AACCCGGTCC CTTTGAACAA GAAGGATCAC
 CCCTCTNGTT TCTTGGGCAC TTGGGCCNGG GAAACTTGTT CTTCCTAGTG
- +1 F N G T I T V T S T L P V N T N D
 6001 TTCAATGGGA CGATCACAGT CACGTCTACN CTGCCAGTGA ACACCAATGA
 AAGTTACCCT GCTAGTGTCA GTGCAGATGN GACGGTCACT TGTGGTTACT
- +1 W I E G E T Y Y C R V T H P H L
 6051 CTGGATCGAG GGCGAGACCT ACTATTGCAG GGTGACCCAC CCGCACCTGC
 GACCTAGCTC CCGCTCTGGA TGATAACGTC CCACTGGGTG GGCGTGGACG
- +1 P K D I V R S I A K A P G K R A P 6101 CCAAGGACAT CGTGCGCTCC ATTGCCAAGG CCCCTGGCAA GCGTGCCCCC GGTTCCTGTA GCACGCGAGG TAACGGTTCC GGGGACCGTT CGCACGGGGG
- +1 P D V Y L F L P P E E E Q G T K D
 6151 CCGGATGTGT ACTTGTTCCT GCCACCGGAG GAGGAGCAGG GGACCAAGGA
 GGCCTACACA TGAACAAGGA CGGTGGCCTC CTCCTCGTCC CCTGGTTCCT
- +1 R V T L T C L I Q N F F P A D I
 6201 CAGAGTCACC CTCACGTGCC TGATCCAGAA CTTCTTCCCC GCGGACATTT
 GTCTCAGTGG GAGTGCACGG ACTAGGTCTT GAAGAAGGGG CGCCTGTAAA
- +1 S V Q W L R N D S P I Q T D Q Y T 6251 CAGTGCAATG GCTGCGAAAC GACAGCCCCA TCCAGACAGA CCAGTACACC GTCACGTTAC CGACGCTTTG CTGTCGGGGT AGGTCTGTCT GGTCATGTGG
- +1 T T G P H K V S G S R P A F F I F 6301 ACCACGGGC CCCACAAGGT CTCGGGCTCC AGGCCTGCCT TCTTCATCTT TGGTGCCCCG GGGTGTTCCA GAGCCCGAGG TCCGGACGGA AGAAGTAGAA
- +1 S R L E V S R V D W E Q K N K F
 6351 CAGCCGCCTG GAGGTTAGCC GGGTGGACTG GGAGCAGAAA AACAAATTCA
 GTCGGCGGAC CTCCAATCGG CCCACCTGAC CCTCGTCTTT TTGTTTAAGT
- +1 T C Q V V H E A L S G S R I L Q K
 6401 CCTGCCAAGT GGTGCATGAG GCGCTGTCCG GCTCTAGGAT CCTCCAGAAA
 GGACGGTTCA CCACGTACTC CGCGACAGGC CGAGATCCTA GGAGGTCTTT

end exon 4

+ 1 W V S K T P G K * 6451 TGGGTGTCCA AAACCCCCGG TAAATGATGC CCACCCTCCT CCCGCCGCCA ACCCACAGGT TTTGGGGGCC ATTTACTACG GGTGGGAGGA GGGCGGCGGT 6501 CCCCCCAGGG CTCCACCTGC TGGGAGGGAG GGGGGCTGGC AAGACCCTCC GGGGGGTCCC GAGGTGGACG ACCCTCCCTC CCCCCGACCG TTCTGGGAGG

polyadenylation signal 6551 ATCTGTCCTT GTCAATAAAC ACTCCAGTGT CTGCTTGGAG CCCTGGGCAC TAGACAGGAA CAGTTATTTG TGAGGTCACA GACGAACCTC GGGACCCGTG

6601 ACCCATTTCT TGGGGGTGGG CAGGGTTGCA GAGCAGGGAT GTCTTGGCAC TGGGTAAAGA ACCCCCACCC GTCCCAACGT CTCGTCCCTA CAGAACCGTG